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INTERFERING RNA MOLECULES

Field of the Invention

The invention provides novel forms of interfering ribonucleic acid molecules having a double-stranded structure. The first strand comprises a first stretch of contiguous nucleotides that is at least partially complementary to a target nucleic acid, and the second strand comprises a second stretch of contiguous nucleotides that is at least partially identical to a target nucleic acid. Methods for using these molecules, for example for inhibiting expression of a target gene, and pharmaceutical compositions, cells and organisms containing these molecules also are provided

Background of the Invention

RNA-mediated interference (RNAi) is a post-transcriptional gene silencing mechanism initiated by double stranded RNA (dsRNA) homologous in sequence to the silenced gene (Fire (1999), Trends Genet 15, 358-63, Tuschl, et al. (1999), Genes Dev 13, 3191-7, Waterhouse, et al. (2001), Nature 411, 834-42, Elbashir, et al. (2001), Nature 411, 494-8, for review see Sharp (2001), Genes Dev 15, 485-90, Barstead (2001), Curr Opin Chem Biol 5, 63-6). RNAi has been used extensively to determine gene function in a number of organisms, including plants (Baulcombe (1999), Curr Opin Plant Biol 2, 109-13), nematodes (Montgomery, et al. (1998), Proc Natl Acad Sci U S A 95, 15502-7), Drosophila (Kennerdell, et al. (1998), Cell 95, 1017-26, Kennerdell, et al. (2000), Nat Biotechnol 18, 896-8). In the nematode C. elegans about one third of the genome has already been subjected to functional analysis by RNAi (Kim (2001), Curr Biol 11, R85-7, Maeda, et al. (2001), Curr Biol 11, 171-6).

Until recently RNAi in mammalian cells was not generally applicable, with the exception of early mouse development (Wianny, et al. (2000), Nat Cell Biol 2, 70-5). The discovery that transfection of duplexes of 21-nt into mammalian cells interfered with gene expression and did not induce a sequence independent interferon-driven anti-viral response usually obtained with long

dsRNA led to new potential application in differentiated mammalian cells (Elbashir et al. (2001), Nature 411, 494-8). Interestingly these small interfering RNAs (siRNAs) resemble the processing products from long dsRNAs suggesting a potential bypassing mechanism in differentiated mammalian cells. The Dicer complex, a member of the RNAse III family, necessary for the initial dsRNA processing has been identified (Bernstein, *et al.* (2001), Nature 409, 363-6, Billy, *et al.* (2001), Proc Natl Acad Sci U S A 98, 14428-33). One of the problems previously encountered when using unmodified ribooligonucleotides was the rapid degradation in cells or even in the serum-containing medium (Wickstrom (1986), J Biochem Biophys Methods 13, 97-102, Cazenave, *et al.* (1987), Nucleic Acids Res 15, 10507-21). It will depend on the particular gene function and assay systems used whether the respective knock down induced by transfected siRNA will be maintained long enough to achieve a phenotypic change.

It is apparent, therefore, that synthetic interfering RNA molecules that are both stable and active in a biochemical environment such as a living cell are greatly to be desired.

Summary of the Invention

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It is therefore an object of the present invention to provide compositions and methods using interfering RNA molecules having enhanced stability.

In accomplishing this object, there has been provided, in accordance with a first aspect of the present invention, a ribonucleic acid comprising a double stranded structure whereby the double- stranded structure comprises a first strand and a second strand, whereby the first strand comprises a first stretch of contiguous nucleotides and whereby said first stretch is at least partially complementary to a target nucleic acid, and the second strand comprises a second stretch of contiguous nucleotides whereby said second stretch is at least partially identical to a target nucleic acid, and whereby the double stranded structure is blunt ended.

In accordance with a second aspect of the present invention there has been provided a ribonucleic acid comprising a double stranded structure whereby the double- stranded structure comprises a first strand and a second strand, whereby the first strand comprises a first stretch of contiguous nucleotides and whereby said first stretch is at least partially complementary to a target nucleic acid, and the second strand comprises a second stretch of contiguous nucleotides, whereby

said second stretch is at least partially identical to a target nucleic acid, whereby the first stretch and/or the second stretch have a length of 18 or 19 nucleotides.

In an embodiment of the ribonucleic acid according to the first aspect of the invention the first stretch and/or the second stretch have a length of 18 or 19 nucleotides.

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In a further embodiment of the ribonucleic acid according to the first aspect of the invention the double stranded structure is blunt ended on both sides of the double strand.

In an alternative embodiment of the ribonucleic acid according to the first aspect of the invention the double stranded structure is blunt ended on the double stranded structure which is defined by the 5'-end of the first strand and the 3'-end of the second strand.

In a further alternative embodiment of the ribonucleic acid according to the first and the second aspect of the invention the double stranded structure is blunt ended on the double stranded structure which is defined by the 3'-end of the first strand and the 5'-end of the second strand.

In accordance with a third aspect of the present invention there has been provided a ribonucleic acid comprising a double stranded structure whereby the double- stranded structure comprises a first strand and a second strand, whereby the first strand comprises a first stretch of contiguous nucleotides and whereby said first stretch is at least partially complementary to a target nucleic acid, and the second strand comprises a second stretch of contiguous nucleotides and whereby said second stretch is at least partially identical to a target nucleic acid, and whereby at least one of the two strands has an overhang of at least one nucleotide at the 5'-end.

In an embodiment of the ribonucleic acid according to the third aspect of the present invention the overhang consists of at least one nucleotide which is selected from the group comprising ribonucleotides and desoxyribonucleotides.

In a more preferred embodiment of the ribonucleic acid according to the third aspect of the present invention the nucleotide has a modification whereby said modification is preferably selected from the group comprising nucleotides being an inverted abasic and nucleotides having an NH₂-modification at the 2'-position.

In a preferred embodiment of the ribonucleic acid according to the third aspect of the present invention at least one of the strands has an overhang of at least one nucleotide at the 3'-end consisting of ribonucleotide or deoxyribonucleotide.

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In another preferred embodiment of the ribonucleic acid according to the third aspect of the present invention the first stretch and/or the second stretch have a length of 18 or 19 nucleotides.

In an embodiment of the ribonucleic acid according to any aspect of the present invention the double-stranded structure has a length of 17 to 21 nucleotides, preferably 18 to 19 nucleotides

In an embodiment of the ribonucleic acid according to the third aspect of the present invention the overhang at the 5'-end is on the second strand.

In a preferred embodiment of the ribonucleic acid according to the third aspect of the present invention the first strand comprises also an overhang, preferably at the 5'-end.

In an embodiment of the ribonucleic acid according to the third aspect of the present invention the 3'-end of the first strand comprises an overhang.

In an alternative embodiment of the ribonucleic acid according to the third aspect of the present invention the overhang at the 5'-end is on the first strand.

In a preferred embodiment thereof the second strand also comprise an overhang, preferably at the 5'-end.

In an embodiment of the ribonucleic acid according to the third aspect of the present invention the 3'-end of the first strand comprises an overhang.

In an embodiment of the ribonucleic acid according to any aspect of the present invention at least one nucleotide of the ribonucleic acid has a modification at the 2'-position and the modification is preferably selected from the group comprising amino, fluoro, methoxy, alkoxy and alkyl.

In accordance with a fourth aspect of the present invention there has been provided a ribonucleic acid comprising a double stranded structure, whereby the double- stranded structure comprises a first strand and a second strand, whereby the first strand comprises a first stretch of contiguous nucleotides and whereby said first stretch is at least partially complementary to a target nucleic acid, and the second strand comprises a second stretch of contiguous nucleotides and whereby said second stretch is at least partially identical to a target nucleic acid,

whereby said first strand and/or said second strand comprises a plurality of groups of modified nucleotides having a modification at the 2'-position whereby within the strand each group of modified nucleotides is flanked on one or both sides by a flanking group of nucleotides whereby the flanking nucleotides forming the flanking group of nucleotides is either an unmodified nucleotide or a nucleotide having a modification different from the modification of the modified nucleotides.

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In an embodiment of the ribonucleic acid according to the fourth aspect of the present invention the ribonucleic acid is the ribonucleic acid according to the first, second or third aspect of the present invention.

In a further embodiment of the ribonucleic acid according to the fourth aspect of the present invention said first strand and/or said second strand comprise said plurality of modified nucleotides.

In another embodiment of the ribonucleic acid according to the fourth aspect of the present invention said first strand comprises said plurality of groups of modified nucleotides.

In yet another embodiment of the ribonucleic acid according to the fourth aspect of the present invention said second strand comprises said plurality of groups of modified nucleotides.

In a preferred embodiment of the ribonucleic acid according to the fourth aspect of the present invention the group of modified nucleotides and/or the group of flanking nucleotides comprises a number of nucleotides whereby the number is selected from the group comprising one nucleotide to 10 nucleotides.

In another embodiment of the ribonucleic acid according to the fourth aspect of the present invention the pattern of modified nucleotides of said first strand is the same as the pattern of modified nucleotides of said second strand.

In a preferred embodiment of the ribonucleic acid according to the fourth aspect of the present invention the pattern of said first strand aligns with the pattern of said second strand.

In an alternative embodiment of the ribonucleic acid according to the fourth aspect of the present invention the pattern of said first strand is shifted by one or more nucleotides relative to the pattern of the second strand.

In an embodiment of the ribonucleic acid according to the fourth aspect of the present invention the modification is selected from the group comprising amino, fluoro, methoxy, alkoxy and alkyl. In another embodiment of the ribonucleic acid according to the fourth aspect of the present invention the double stranded structure is blunt ended.

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In a preferred embodiment of the ribonucleic acid according to the fourth aspect of the present invention the double stranded structure is blunt ended on both sides.

In another embodiment of the ribonucleic acid according to the fourth aspect of the present invention the double stranded structure is blunt ended on the double stranded structure's side which is defined by the 5'-end of the first strand and the 3'-end of the second strand.

In still another embodiment of the ribonucleic acid according to the fourth aspect of the present invention the double stranded structure is blunt ended on the double stranded structure's side which is defined by at the 3'-end of the first strand and the 5'-end of the second strand.

In another embodiment of the ribonucleic acid according to the fourth aspect of the present invention at least one of the two strands has an overhang of at least one nucleotide at the 5'-end.

In a preferred embodiment of the ribonucleic acid according to the fourth aspect of the present invention the overhang consists of at least one desoxyribonucleotide.

In a further embodiment of the ribonucleic acid according to the fourth aspect of the present invention at least one of the strands has an overhang of at least one nucleotide at the 3'-end.

In an embodiment of the ribonucleic acid according to any of the aspects of the present invention the length of the double-stranded structure has a length from about 17 to 21 and more preferably 18 or 19 bases

In another embodiment of the ribonucleic acid according to any of the aspects of the present invention the length of said first strand and/or the length of said second strand is independently from each other selected from the group comprising the ranges of from about 15 to about 23 bases, 17 to 21 bases and 18 or 19 bases.

In a preferred embodiment of the ribonucleic acid according to any of the aspects of the present invention the complementarity between said first strand and the target nucleic acid is perfect.

In an embodiment of the ribonucleic acid according to any of the aspects of the present invention the duplex formed between the first strand and the target nucleic acid comprises at least 15 nucleotides wherein there is one mismatch or two mismatches between said first strand and the target nucleic acid forming said double-stranded structure.

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In an embodiment of the ribonucleic acid according to any of the aspects of the present invention, wherein both the first strand and the second strand each comprise at least one group of modified nucleotides and at least one flanking group of nucleotides, whereby each group of modified nucleotides comprises at least one nucleotide and whereby each flanking group of nucleotides comprising at least one nucleotide; with each group of modified nucleotides of the first strand being aligned with a flanking group of nucleotides on the second strand, whereby the most terminal 5' nucleotide of the first strand is a nucleotide of the group of modified nucleotides, and the most terminal 3' nucleotide of the second strand is a nucleotide of the flanking group of nucleotides.

In a preferred embodiment of the ribonucleic acid according to the fourth aspect, wherein each group of modified nucleotides consists of a single nucleotide and/or each flanking group of nucleotides consists of a single nucleotide.

In a further embodiment of the ribonucleic acid according to the fourth aspect, wherein on the first strand the nucleotide forming the flanking group of nucleotides is an unmodified nucleotide which is arranged in a 3' direction relative to the nucleotide forming the group of modified nucleotides, and wherein on the second strand the nucleotide forming the group of modified nucleotides is a modified nucleotide which is arranged in 5' direction relative to the nucleotide forming the flanking group of nucleotides.

In another embodiment of the ribonucleic acid according to the fourth aspect, wherein the first strand comprises eight to twelve, preferably nine to eleven, groups of modified nucleotides, and wherein the second strand comprises seven to eleven, preferably eight to ten, groups of modified nucleotides.

In a preferred embodiment of the ribonucleic acid according to any of the aspects of the present invention the target gene is selected from the group comprising structural genes, housekeeping genes, transcription factors, motility factors, cell cycle factors, cell cycle inhibitors, enzymes, growth factors, cytokines and tumor suppressors.

In a further embodiment of the ribonucleic acid according to any of the aspects of the present invention the first strand and the second strand are linked by a loop structure.

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In a preferred embodiment of the ribonucleic acid according to any of the aspects of the present invention the loop structure is comprised of a non-nucleic acid polymer.

In a preferred embodiment thereof the non-nucleic acid polymer is polyethylene glycol.

In an alternative embodiment thereof the loop structure is comprised of a nucleic acid.

In an embodiment of the ribonucleic acid according to any of the aspects of the present invention the 5'-terminus of the first strand is linked to the 3'-terminus of the second strand.

In a further embodiment of the ribonucleic acid according to any of the aspects of the present invention the 3'-end of the first strand is linked to the 5'-terminus of the second strand.

In accordance with a fifth aspect of the present invention there have been provided methods of using a he ribonucleic acid according to any of the aspects of the present invention for target validation.

In accordance with a sixth aspect of the present invention there have been provided medicaments and pharmaceutical compositions containing a ribonucleic acid according to any of the aspects of the present invention, and methods of making such medicaments and compositions.

In a preferred embodiment of the use according to the sixth aspect of the present invention methods are provided for the treatment of a disease or of a condition which is selected from the group comprising glioblastoma, prostate cancer, breast cancer, lung cancer, liver cancer, colon cancer, pancreatic cancer and leukaemia, diabetes, obesity, cardiovascular diseases, and metabolic diseases.

In accordance with a seventh aspect of the present invention there has been provided a cell, for example a knockdown cell, containing a ribonucleic acid according to any of the aspects of the present invention.

In accordance with an eighth aspect of the present invention there has been provided an organism, for example a knockdown organism, containing a ribonucleic acid according to any of the aspects of the present invention.

In accordance with a ninth aspect of the present invention there has been provided a composition containing a ribonucleic acid according to any of the aspects of the present invention.

In accordance with a tenth aspect of the present invention there has been provided a pharmaceutical composition containing a ribonucleic acid according to any of the aspects of the present invention, and a pharmaceutically acceptable carrier.

In accordance with an eleventh aspect of the present invention there has been provided a method for inhibiting the expression of a target gene in a cell or derivative thereof comprising introducing a ribonucleic acid according to any of the aspects of the present invention into a cell in an amount sufficient to inhibit expression of the target gene, wherein the target gene is the target gene of the a ribonucleic acid according to any of the aspects of the present invention.

Brief Description of the Drawings

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Fig. 1 shows a schematic illustration defining the terminology as used herein. The upper of the two strands is the first strand and the antisense strand of the targeted nucleic acid such as mRNA. The second strand is the one which essentially corresponds in its sequence to the targeted nucleic acid and thus forms the sense strand. Both, the first strand and second strand form a double-stranded structure, typically through Watson Crick base pairing.

Fig. 2 illustrates some embodiments of the ribonucleic acid molecules of the present invention with patterns of modified and unmodified groups of nucleotides which are also referred to herein as a pattern of modification. The modified groups of nucleotides are also referred to herein as a group of modified nucleotides. The unmodified nucleotides or unmodified groups of nucleotides referred to as flanking group(s) of nucleotides herein, as used herein may also have one or several of the modification(s) as disclosed herein which, however, is/are different from the modification of the nucleotides forming the group(s) of modified nucleotides. In Fig. 2A the modified and unmodified groups of nucleotides, *i.e.* the groups of modified nucleotides and the flanking groups of nucleotides on both the first stretch and the second stretch are located on corresponding parts of the stretches and are thus aligned to each other (groups of modified nucleotides on the first strand aligned with groups of modified nucleotides on the second strand and flanking groups of nucleotides on the first

strand aligned with flanking group of nucleotides on the second strand), whereas in Fig. 2B the pattern realised on the first strand is also realised on the second strand, however, with a phase shift such that the modified group of nucleotides of the first stretch is base pairing with an unmodified group of nucleotides of the second stretch and vice versa so that a group of modified nucleotides on the first strand aligns with a flanking group of nucleotides on the second strand. In Fig. 2C a further possibility of arranging the modified and unmodified groups of nucleotides is realised. It is also within the present invention that the pattern of the first stretch is independent from the pattern of the second stretch and that both patterns partially overlap in terms of relative position to each other in the double-stranded structure defined by base pairing. In a further embodiment the extent of this overlapping can vary over the length of the stretch(es) and strand(s), respectively.

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Fig. 3 shows the result of a knockdown experiment using RNAi molecules with different end protection groups. More particularly Fig. 3A shows that the various forms of end protected RNAi molecules are functional on the knockdown of PTEN mRNA.

Fig. 3B shows the sequence of the different RNAi molecules used in the experiment the result of which is depicted in Fig. 3A. Fig. 3C shows the result of an immunoblot analysis of PTEN protein after treatment with modified RNAi molecules in comparison to PTEN specific antisense constructs.

Fig. 4 shows that the 3' overhang of RNAi molecules is not important for RNA interference. More particularly, Fig. 4A shows a dose response curve of different RNAi molecules and Fig. 4B shows the sequence of the RNAi molecules used in the experiment the result of which is shown in Fig. 4A.

Fig. 5 shows that duplex length of the RNAi molecules has to be at least 18-19 nucleotides. More particularly, Fig. 5B shows the sequence of the PTEN specific RNAi molecules used in the experiment the result of which is depicted in Fig. 5A as dose response curve.

Fig. 6 shows that four terminal mismatched nucleotides in RNAi molecules with a length of 19 nucleotides are still functional in mediating Akt1 knockdown. More particularly, Fig. 6B shows the sequence of the RNAi molecules used in the experiment the result of which is depicted in Fig. 6A.

Fig. 7 shows further results on duplex length requirements and tolerance for mutation in siRNAs. More particularly, Fig. 7A shows the various constructs used (left panel) and the respective impact on inhibition of Akt1 mRNA expression in HeLa cells relative to the expression of p110α used in the indicated amounts of siRNA molecules (right panel). The nucleotide changes in the mismatch siRNA molecules are indicated by arrows; the 3' desoxynucleotides, if any, are indicated in capital letters. Fig. 7B shows the various PTEN specific siRNAs (left panel), the inhibition of PTEN mRNA expression in HeLA cells expressed as ratio PTEN/p110α, at various amounts of siRNA (middle panel) and Fig. 7C a Western Blot analysis depicting the inhibition of PTEN protein expression using PTEN specific siRNA (30nM) and respective mismatch siRNA after 48 and 96 hours, respectively, with p100α being used as loading control.

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Fig. 8 shows the result of studies on the stability in serum conferred to RNAi molecules by 2'-O-methylation and that end modifications have no beneficial effects on RNAi stability. More particularly, Fig. 8A shows the result of a gel electrophoresis of the various RNAi molecules depicted in Fig. 8B being subject to incubation with fetal calf serum.

Fig. 9 shows that an amino end modification results in loss of activity. Fig. 9B shows the particular RNAi molecules used in the experiments the result of which is shown in Fig. 9A expressed as PTEN/p110α expression level ratio. Fig. 9C shows the design principles which may be deduced from the results depicted in Fig. 9A. As used in Fig. 9C the term functional means functionally active in the particular assay system as described in the example and "not functional" means not functionally active in said system.

Fig. 10 shows that 2'-O-Alkyl (methyl) modifications stabilize RNAi molecules but also result in reduction of their activity. More particularly, Fig. 10C shows the sequence of the RNAi molecules used in the experiment the result of which is depicted as a dose response curve in Fig. 10A. Fig. 10B shows the result of a gel electrophoresis of the various RNAi molecules depicted in Fig. 10C being subject to a two hour incubation in fetal calf serum.

Fig. 11 shows the result of an experiment on the efficacy of RNAi molecules with blocks of 2'-O-methyl modifications with Fig. 11A graphically depicting the results of said experiments as a dose response curve and with Fig. 11C showing the sequences of the particular RNAi molecules used in said

experiments. Fig. 11B shows the result of a gel electrophoresis of the various RNAi molecules depicted in Fig. 11C being subject to a two hour incubation in fetal calf serum.

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Fig. 12 shows that alternating 2'-O-methyl modification result in activity of the modified RNAi molecules compared to unmodified forms. More particularly, Fig. 12B shows the sequence of the RNAi molecules used in this experiment the result of which is depicted in Fig. 12A. Fig. 12C shows the stability of said RNAi molecules following incubation in serum for two hours, whereas Fig. 12D shows an immunoblot for PTEN protein upon application of different RNAi molecules to HeLa cells. As may be taken therefrom RNAi molecules with alternating modifications are stabilized against endonuclease degradation and active in mediating a PTEN protein knock down.

Fig. 13 shows the result of a Western Blot analysis to determine the time course of PTEN protein knock down. Cells were continuously transfected with 2'-O-Methyl modified versus unmodified RNAi molecules using cationic lipids for 72 h. Protein lysates were prepared and analysed by immunoblot after 48 and 120 h. For the 96 h and 120 h timepoints the cells were split, replated and incubated in the absence of RNAi molecules for an additional 24 and 48 h.

Fig. 14 shows a Western Blot depicting the protein knock down of PTEN being persistent using alternating modified RNAi molecules versus unmodified RNAi molecules. Transfections were performed for only 5 h and new medium without transfection reagents was added. Lysates were analysed by immunoblot 72h and 96h post transfection with the indicated RNAi molecules.

Fig. 15 shows that siRNA molecules with distinct 2'-O-methyl ribonucleotides modifications show increased stability in serum and mediate protein knock-down in HeLa cells. More particularly, Fig. 15A indicates the various siRNA molecule constructs used (left panel), whereby 2'-O-methyl ribonucleotides modifications are underlined and indicated by bold letters in the sequence. Inhibition of PTEN mRNA expression in HeLa cells transfected with the indicated amounts of modified siRNA molecules is expressed as ratio PTEN/p110 α and indicated on the right panel. Fig. 15B shows on the left panel the various siRNA constructs used and on the right panel a PAA gel electrophoresis of modified and unmodified siRNA molecules after incubation in serum; the various constructs with 2'-O-methyl ribonucleotides are indicated by

underlining and bold printing. Fig. 15C shows an SDS-PAGE based immunoblot illustrating the inhibition of PTEN protein expression using various of the siRNA constructs (30 nM) as depicted in Fig. 15A and 15B, respectively. Again, p110 α is used as loading control. Finally, Fig. 15D is an immunoblot indicating a prolonged protein knock-down, i.e. the inhibition of PTEN protein expression, upon administration of siRNA molecules (30 nM) with distinct 2'-O-methylribonucleotides modifications after 48 and 128 hours. As in Fig. 15C, p 110 α is used as loading control.

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Fig. 16 shows that siRNA molecules with distinct 2'-O-10 methylribonucleotides modifications which are specific for Akt1 and p110ß mRNA show increased stability in serum and mediate protein knock-down in HeLa cells. More particularly, Fig. 16A indicates on the left panel the various constructs used whereby again 2'-O-methylribonucleotides are underlined and printed in bold. The integrity of the indicated siRNA molecules after incubation in 15 serum is shown in the right panel. Fig. 16B shows an immunoblot of Akt1, Akt2 and Akt phosphorylation and p110 being used as a loading control upon transfection of the cells with the indicated siRNAs (30 mM). Fig. 16C shows various p110β specific siRNA constructs (left panel) with the 2'-O-methyl modifications being underlined and printed in bold, and the result of an 20 immunoblot analysis (right panel) of the inhibition of the phosphorylation of the downstream kinase Akt1 by said siRNA constructs. p110α has been used as a loading control.

Fig. 17 shows the efficacy of various RNAi molecules with hairpin structures as dose response curve while Fig. 17B shows the structure of the RNAi molecules the result of which is depicted in Fig. 17A. Synthetic siRNAs with different loops are functional in reducing the p110β, Akt1 and Akt 2 expression. (14A) Inhibition of p110β mRNA expression in siRNA transfected HeLa cells. Samples were analyzed in parallel for the level of p110β mRNA expression 24h after transfection of the indicated siRNAs. The transfected bimolecular siRNAs (21mer with 3' TT overhangs, molecule 1AB) or the monomolecular siRNAs with loop structures are schematically shown. Note that the position of the loops (HIV derived pA-loop; (A)₁₂-loop) relative to the antisense sequence is reversed in 3A, 4A relative to 3B, 4B. The 2AB siRNA molecule contains 6 mismatches in the 21mer duplex and serves as a negative control together with the untreated sample.

RNA was prepared und subjected to real time RT-PCR (Taqman) analysis. p110 β mRNA levels are shown relative to the mRNA levels of p110 α , which serve as an internal reference. Each bar represents triplicate transfections (\pm standard deviation). HeLa cells were transfected at 50% confluency (2500 cells per 96 well) with siRNAs at the indicated concentrations in growth medium.

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Fig. 18 shows the efficacy of various RNAi molecules with intermolecular and intramolecular loop structures as dose response curves. (18A) Inhibition of Akt1 mRNA expression in siRNA transfected HeLa cells. Samples were analysed in parallel for the level of Akt1 and Akt2 mRNA expression 24h after transfection of the indicated siRNAs. The different loops (A-loops; GAGA-loop and a polyethyleneglycol (PEG)-linker) and their putative secondary structure are shown schematically. The siRNA molecule 9A is specific for Akt2 and serves as a negative control. Note that 10A and 10B do not contain self-complementary sequences and are transfected in combination. Akt1 mRNA levels is shown relative to the mRNA levels of p110β, which served as internal control. (18B) Inhibition of Akt2 mRNA expression in HeLa cells transfected with the indicated siRNA molecules. Akt2 mRNA levels is shown relative to the mRNA levels of p110β. The Akt1 specific molecule 7A serves here as a negative control.

Fig. 18 C shows a Western Blot analysis on Akt protein depicting the functionality of synthetic siRNAs with different loops in specifically reducing the Akt1 and Akt 2 expression. Inhibition of Akt1 and Akt2 protein expression were analysed by immunoblot. The cells were harvested 48h after transfection of the indicated hairpin siRNAs (20nM). Cell extracts were separated by SDS-PAGE and analysed by immunoblotting using anti-p110 antibody, anti Akt 1/2. Similar results were obtained with an antibody specific for the phosphorylated form of Akt1. The positions of p110α, another catalytic subunit of PI 3-kinase, which was used as a loading control, and of Akt1, Akt2 and phosphorylated Akt (P*-Akt) are indicated on the left.

Fig. 19 shows an NH₂ modification, also referred to herein as amino modification, which may be present at either the 3'-OH terminal nucleotide or the 5' terminal nucleotide. The amino group is attached to the phosphate which in turn is attached to the OH group of the sugar moiety, through an alkyl group comprising an alkyl chain of 1 to 8, preferably 6 C atoms, whereby the second C atom close to the phosphate group has a CH₂OH group attached thereto. As an

alternative the linker may be formed by an ether whereby the ether is comprised of two alcohols whereby one alcohol is an amino alcohol and the other is a dialcohol with one alcohol group involved in the formation of the ether group and the other one being an OH group located at either of the C atoms, preferably at the second C atom relative to the phosphate group. Detailed Description

Detailed Description

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The present inventors have surprisingly found that small interfering RNAs can be designed that are both highly specific and active as well as stable under the reaction conditions typically encountered in biological systems such as biochemical assays or cellular environments. The various interfering RNAs previously described by Tuschl *et al.* (see, for example, WO01/75164) provide for a length of 21 to 23 nucleotides and a modification at the 3' end of the double-stranded RNA. The present inventors found that stability problems of interfering RNA, including small interfering RNA (siRNA) which is generally referred to herein in the following as RNAi, actually result from attack by endonucleases rather than exonucleases as previously thought. This surprising observation permitted the present inventors to develop the methods and compositions that are the subject of the present invention.

Structure of RNAi molecules

The present invention provides new forms of interfering RNA. RNAi consists of a ribonucleic acid comprising a double-stranded structure, formed by a first strand and a second strand. The first strand comprises a stretch of contiguous nucleotides (the first stretch of contiguous nucleotides) that is at least partially complementary to a target nucleic acid. The second strand also comprises a stretch of contiguous nucleotides where the second stretch is at least partially identical to a target nucleic acid. The very basic structure of this ribonucleic acid is schematically shown in Fig. 1. The first strand and said second strand may be hybridized to each other to form a double-stranded structure. The hybridization typically occurs by Watson Crick base pairing.

The ribonucleic acids of the invention are, however, not necessarily limited in length to such a double-stranded structure. For example, further nucleotides may added to each strand and/or to each end of any of the strands forming the RNAi. Depending on the particular sequence of the first stretch and the second stretch, the hybridization or base pairing is not necessarily complete or

perfect, which means that the first stretch and the second stretch are not 100 % base paired due to mismatches. One or more mismatches may also be present within the duplex. Such mismatches have no effect on the RNAi activity if placed outside a stretch of preferably 15, 16 or 17 matching nucleotides. If mismatches are placed to yield only 15 or less contiguous matching nucleotides, the RNAi molecule typically shows a reduced activity in down regulating mRNA for a given target compared to a 17 matching nucleotide duplex.

The first stretch of contiguous nucleotides of the first strand is essentially complementary to a target nucleic acid, more advantageously to a part of the target nucleic acid. The term "complementary" as used herein preferably means that the nucleotide sequence of the first strand hybridizes to a nucleic acid sequence of a target nucleic acid sequence or a part thereof. Typically, the target nucleic acid sequence or target nucleic acid is, in accordance with the mode of action of interfering ribonucleic acids, a single stranded RNA, more preferably an mRNA. Such hybridization occurs most likely through Watson Crick base pairing but, however, is not necessarily limited thereto. The extent to which the first strand and more particularly the first stretch of contiguous nucleotides of the first strand is complementary to a target nucleic acid sequence can be as high as 100% and be as little as 80%, advantageously 80-100%, more advantageously 85-100%, most advantageously 90-100%. Optimum complementarity seems to be 95-100%. Complementarity in this sense means that the aforementioned range of nucleotides, such as, e.g., 80%-100%, depending on the particular range, of the nucleotides are perfect by Watson Crick base pairing.

Complementarity of the strands

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It is shown in one aspect of the present invention that the complementarity between the first stretch of nucleotides and the target RNA has to be 18-19 nucleotides; stretches of as little as 17 nucleotides even with two sequence specific overhangs are not functional in mediating RNAi. Accordingly, in a duplex having a length of 19 nucleotides or base pairs a minimum complementarity of 17 nucleotides or nucleotide base pairs is acceptable, allowing for a mismatch of two nucleotides. In the case of a duplex consisting of 20 nucleotides or base pairs a complementarity of 17 nucleotides or nucleotide base pairs is allowable and functionally active. The same principle applies to a duplex of 21 nucleotides or base pairs with a total of 17 complementary nucleotides or

base pairs. The extent of complementarity required for a length of a duplex, i.e. of a double stranded structure, can also be based on the melting temperature of the complex formed by either the double stranded structure as described herein or by the complex of the first stretch of the first strand and the target nucleic acid.

The skilled artisan will understand that all of the ribonucleic acids of the present invention are suitable to cause or being involved in methods of RNA mediated interference such as those described, for example, in WO 99/32619, WO 00/44895 and WO 01/75164.

Length of the RNAi molecules

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The first strategy by which an interfering ribonucleic acid molecule may be designed according to the present invention is to have an optimum length of 18 or 19 nucleotides of the stretch that is complementary to the target nucleic acid. It is also within the scope of the present invention that this optimum length of 18 or 19 nucleotides is the length of the double stranded structure in the RNAi used.

This length requirement clearly differs from that described in, for example, WO 01/75164. It is within the scope of the present invention that any further design, both according to the present invention and as previously described by others, can be realised in connection with an interfering ribonucleic acid having such length characteristics, i.e. a length of 18 or 19 nucleotides.

End modification of the RNAi molecules

The second strategy for the design of an interfering ribonucleic acid molecule is to have a free 5' hydroxyl group, (also referred to herein as a free 5' OH-group) at the terminus of the first strand. A free 5' OH-group means that the most terminal nucleotide forming the first strand is present and is thus not modified, particularly not by an end modification. Typically, the terminal 5'-hydroxy group of the second strand, respectively, is also present in an unmodified manner. In a more preferred embodiment, the 3'-end of the first strand and first stretch, respectively, also is unmodified so as to present a free OH-group (also referred to herein as a free 3'OH-group), whereby the design of the 5' terminal nucleotide is the one of any of the embodiments described above. Preferably such a free OH-group also is present at the 3'-end of the second strand and second stretch, respectively. In other embodiments of the ribonucleic acid molecules as described previously according to the present invention the 3'-end of the first

strand and first stretch, respectively, and/or the 3'-end of the second strand and second stretch, respectively, may have an end modification at the 3' end.

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As used herein the terms free 5'OH-group and 3'OH-group also indicate that the respective 5' and 3' terminal nucleotides of the polynucleotide present an OH-group. Such OH-group may stem from either the sugar moiety of the nucleotide, more preferably from the 5' position in the case of the 5'OH-group and from the 3' position in the case of the 3'OH-group, or from a phosphate group attached to the sugar moiety of the respective terminal nucleotide. The phosphate group may in principle be attached to any OH-group of the sugar moiety of the nucleotide. Preferably, the phosphate group is attached to the 5'OH-group of the sugar moiety in the case of the free 5'OH-group and/or to the 3'OH-group of the sugar moiety in the case of the free 3'OH-group still providing what is referred to herein as free 5' or 3' OH-group.

The term "end modification" as used herein in connection with any strategy for the design of RNAi or any embodiment of RNAi disclosed herein, means a chemical entity added to the most 5' or 3' nucleotide of the first and/or second strand. Examples of such end modifications include, but are not limited to, inverted (deoxy) abasics, amino, fluoro, chloro, bromo, CN, CF, methoxy, imidazole, carboxylate, thioate, C₁ to C₁₀ lower alkyl, substituted lower alkyl, alkaryl or aralkyl, OCF₃, OCN, O-, S-, or N-alkyl; O-, S-, or N-alkenyl; SOCH₃; SO₂CH₃; ONO₂; NO₂, N₃; heterocycloalkyl; heterocycloalkaryl; aminoalkylamino; polyalkylamino or substituted silyl, as, among others, described in European patents EP 0 586 520 B1 or EP 0 618 925 B1.

As used herein, alkyl or any term comprising "alkyl" means any carbon atom chain comprising 1 to 12, preferably 1 to 6 and more, preferably 1 to 2 C atoms.

A further end modification is a biotin group, which may preferably be attached to either the most 5' or the most 3' nucleotide of the first and/or second strand, or to both ends. In a more preferred embodiment the biotin group is coupled to a polypeptide or a protein. It is also within the scope of the present invention that the polypeptide or protein is attached through any of the end modifications described herein. The polypeptide or protein may confer further characteristics to the nucleic acid molecules of the invention. For example, the polypeptide or protein may act as a ligand to another molecule. If the other

molecule is a receptor the receptor's function and activity may be activated by the binding ligand. The receptor may show an internalization activity which provides for effective transfection of the ligand bound inventive nucleic acid molecules. An example of a ligand that may be coupled to the nucleic acid molecules of the invention is VEGF and the corresponding receptor is the VEGF receptor.

Various possible embodiments of the RNAi of the present invention having different kinds of end modification(s) are presented in the Table 1 below:

Table 1: Various embodiments of the interfering ribonucleic acid according to the present invention

		1 st strand/1 st stretch	2 nd strand/ 2nd
			stretch
1.)	5'-end	free OH	free OH
	3'-end	free OH	free OH
2.)	5'-end	free OH	free OH
	3'-end	end modification	end modification
3.)	5'-end	free OH	free OH
	3'-end	free OH	end modification
4.)	5'-end	free OH	free OH
	3'-end	end modification	free OH
5.)	5'-end	free OH	end modification
	3'-end	free OH	free OH
6.)	5'-end	free OH	end modification
	3'-end	end modification	free OH
7.)	5'-end	free OH	end modification
	3'-end	free OH	end modification
8.)	5'-end	free OH	end modification
•	3'-end	end modification	end modification

The various end modifications as disclosed herein are preferably located at the ribose moiety of a nucleotide of the ribonucleic acid. More particularly, the end modification may be attached to or replace any of the OH-groups of the ribose moiety, including but not limited to the 2'OH, 3'OH and 5'OH position, provided that the nucleotide thus modified is a terminal nucleotide. Inverted abasics are nucleotides, either desoxyribonucleotides or ribonucleotides which do not have a

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nucleobase moiety. This kind of compound is, among others, described in Sternberger et al.(2002), Antisense. Nucl. Ac. Drug Dev. 12:143.

Any of the aforementioned end modifications may be used in connection with the various embodiments of RNAi depicted in table 1. It is particularly advantageous to inactivate the sense strand of any of the RNAi forms or embodiments disclosed herein, preferably via an end modification, and more preferably a 5' end modification. The advantage of this strategy arises from the inactivation of the sense strand which corresponds to the second strand of the ribonucleic acids described herein, which might otherwise interfere with an unrelated single-stranded RNA in the cell. Thus the expression and more particularly the translation pattern of the transcriptome of a cell is more specifically influenced. This effect is also referred to as an off-target effect. Referring to Table 1 those embodiments depicted as embodiments 7 and 8 are particularly advantageous in the above sense as the modification results in an inactivation of the - target unspecific - part of the RNAi (which is the second strand) thus reducing any unspecific interaction of the second strand with singlestranded RNA in a cellular or similar system where the RNAi according to the present invention is going to be used to knock down specific ribonucleic acids and proteins, respectively.

20 Blunt-ended RNAi molecules

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A third strategy provided by the present invention involves a ribonucleic acid comprising a double-stranded structure having a first strand and a second strand, where the first strand comprises a first stretch of contiguous nucleotides and where the first stretch is at least partially complementary to a target nucleic acid, and the second strand comprises a second stretch of contiguous nucleotides where the second stretch is at least partially identical to a target nucleic acid, where the double-stranded structure is blunt-ended. As used herein the term double-stranded structure also is referred to as duplex. This design of RNAi clearly differs from, *e.g.*, that described by Tuschl *et al.* in WO 01/75164, which contains a 3'-overhang. As used herein the term overhang refers to a double-stranded structure where at least one end of one strand is longer than the corresponding end of the other strand ("the counter strand") forming the double-stranded structure. Preferably, the first stretch is identical to the first strand and the second stretch is identical to the second strand.

It also is advantageous to employ a combination of the design principles described above, *i.e.* to prepare RNAi molecules that are blunt-ended and that have an end modification of either the first or the second strand or both. In other words, it is within the scope of the present invention to have blunt-ended RNAi carrying any end modification scheme as depicted in Table 1.

RNAi molecules having a 5' overhang

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The fourth strategy of the present invention is to have an overhang at the 5'-end of the ribonucleic acid. More particularly, such overhang may in principle be present at either or both the first strand and second strand of the ribonucleic acid according to the present invention. The length of the overhang may be as little as one nucleotide and as long as 2 to 8 nucleotides, preferably 2, 4, 6 or 8 nucleotides. It is within the present invention that the 5' overhang may be located on the first strand and/or the second strand of the ribonucleic acid. The nucleotide(s) forming the overhang may be (a) desoxyribonucleotide(s), (a) ribonucleotide(s) or a continuation thereof.

The overhang preferably comprises at least one desoxyribonucleotide, whereby said one desoxyribonucleotide is preferably the most 5'-terminal one. It is within the present invention that the 3'-end of the respective counter-strand of the inventive ribonucleic acid does not have an overhang, more preferably not a desoxyribonucleotide overhang. Here again, any of the inventive ribonucleic acids may comprise an end modification scheme as outlined in connection with table 1 and/or and end modification as outlined herein.

RNAi molecules having modified nucleotides

A fifth strategy for the design of interfering ribonucleic acids subject to the present application resides in the formation of a certain pattern of modified nucleotides on at least one of the strands and more particularly on one of the stretches of contiguous nucleotides of the ribonucleic acid(s) according to the present invention. The type of modification of the nucleotides may be the same as that discussed in connection with the other strategies for designing interfering RNA disclosed herein. Advantageously, the modification is an end modification as described herein, such as, e.g., inverted abasics, methoxy, or amino and the like at the ribose moiety of at least one nucleotide forming the ribonucleotide acids according to the present application. The modification of the nucleotides may be any form of modification described herein, more particularly the kind of

modification as described herein as end modification, except that the so-called end modification is not necessarily located at terminal nucleotides. Rather the modification may occur at a non-terminal nucleotide. Under such conditions the modification is preferably attached to the ribose moiety of the - to be - modified nucleotide and even more preferably to the 2' position of the ribose moiety.

Combinations of design elements

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It is also within the present invention that any ribonucleic acid designed according to this strategy may also have the features conferred to a ribonucleic acid according to the present application by any of the other design strategies disclosed herein. Accordingly, the interfering ribonucleic acid having a pattern of modified nucleotides may have an end modification, an end modification scheme, may be blunt ended or may have a 5' overhang or any combination of two or more of these elements or characteristics.

Apart from the aforementioned modifications which may be presented either as end modifications or as modification pattern, the ribonucleic acid backbone as such may be further modified by forming different links between the nucleotides. Examples of such different links are described, among others, in EP 0 586 520 B1 and EP 0 618 925 B1. Of particular interest here are internal modification(s) of the ribonucleic acid backbone which have been shown to confer higher nuclease resistance of ribooligonucleotides. In a preferred embodiment the modification of the modified nucleotide is a methylation of the 2'-OH-group of the ribose moiety of the nucleotide to form a methoxy group.

In a preferred embodiment, both strands, and more particularly both the first stretch and the second stretch show this kind of modification of the nucleotides forming said strands and stretches, respectively. However, it is also within the present invention that either the first strand and first stretch, respectively, or the second strand and second stretch, respectively, show this particular pattern of modification of the nucleotides. As used herein, the term group of modified nucleotides or flanking group of nucleotides may comprise or represent one or more nucleotides.

A pattern of modification of the nucleotides in a contiguous stretch of nucleotides may be realised such that the modification is contained within a single nucleotide or group of nucleotides that are covalently linked to each other via standard phosphodiester bonds or, at least partially, through phosphorothioate

bonds. In the event that such a modified nucleotide or group of modified nucleotides does not form the 5'-end or 3'-end of the stretch, a flanking nucleotide or group of nucleotides is arrayed on both sides of the modified nucleotide or group, where the flanking nucleotide or group either is unmodified or does not have the same modification of the preceding nucleotide or group of nucleotides. The flanking nucleotide or group of nucleotides may, however, have a different modification. This sequence of modified nucleotide or group of modified nucleotides, respectively, and unmodified or differently modified nucleotide or group of unmodified or differently modified nucleotides may be repeated one or more times. Preferably, the sequence is repeated more than one time. The pattern of modification is discussed in more detail below, generally referring to a group of modified nucleotides or a group of unmodified nucleotides whereby each of said group may actually comprise as little as a single nucleotide. The term "unmodified nucleotide" as used herein means either not having any of the aforementioned modifications at the nucleotide forming the respective nucleotide or group of nucleotides, or having a modification which is different from the one of the modified nucleotide and group of nucleotides, respectively.

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It is also within the present invention that the modification of the unmodified nucleotide(s) wherein such unmodified nucleotide(s) is/are actually modified in a way different from the modification of the modified nucleotide(s), can be the same or even different for the various nucleotides forming said unmodified nucleotides or for the various flanking groups of nucleotides. That is, a group of modified nucleotides may contain two or more different modified nucleotides within a single group.

The pattern of modified and unmodified nucleotides may be such that the 5'-terminal nucleotide of the strand or of the stretch starts with a modified group of nucleotides or starts with an unmodified group of nucleotides. However, in an alternative embodiment it is also possible that the 5'-terminal nucleotide is formed by an unmodified group of nucleotides.

This kind of pattern may be realised either on the first stretch or the second stretch of the interfering RNA or on both. A 5' phosphate on the target-complementary strand of the siRNA duplex is required for siRNA function, suggesting that cells check the authenticity of siRNAs through a free 5' OH

(which can be phosphorylated) and allow only such bona fide siRNAs to direct target RNA destruction (Nykanen, et al. (2001), Cell 107, 309-21).

Preferably, the first stretch shows a pattern of modified and unmodified groups of nucleotides, i.e. of group(s) of modified nucleotides and flanking group(s) of nucleotides, whereas the second stretch does not show this kind of pattern. This may be useful insofar as the first stretch is actually the more important one for the target-specific degradation process underlying the interference phenomenon of RNA so that for specificity reasons the second stretch can be chemically modified so it is not functional in mediating RNA interference.

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However, it is also within the present invention that both the first stretch and the second stretch have this kind of pattern. Preferably, the pattern of modification and non-modification is the same for both the first stretch and the second stretch.

In a particular embodiment, the modified nucleotides or groups of modified nucleotides of one strand of the molecule are complementary in position to the modified nucleotides or groups of nucleotides of the other strand. This possibility is schematically depicted in Fig. 2A. In an alternative embodiment, there is a phase shift between the patterns of modification of the first stretch and first strand, respectively, relative to the pattern of modification of the second stretch and second strand, respectively. Preferably, the shift is such that the modified group of nucleotides of the first strand corresponds to the unmodified group of nucleotides of the second strand and vice versa. This possibility is illustrated schematically in Fig. 2B. In another embodiment, there is a partial shift of the pattern of modification to that the modified groups overlap as illustrated in Fig. 2C. The groups of modified nucleotides in any given strand may optionally be the same length, but may be of different lengths. Similarly, The groups of unmodified nucleotides in any given strand may optionally be the same length, or of different lengths.

In a preferred embodiment the second (penultimate) nucleotide at the terminus of the strand and stretch, respectively, is an unmodified nucleotide or the beginning of group of unmodified nucleotides. Preferably, this unmodified nucleotide or unmodified group of nucleotides is located at the 5'-end of the first and second strand, respectively, and even more preferably at the terminus of the first strand. In a further preferred embodiment the unmodified nucleotide or

unmodified group of nucleotide is located at the 5'-end of the first strand and first stretch, respectively. In a preferred embodiment the pattern consists of alternating single modified and unmodified nucleotides.

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In a particular embodiment of this aspect of the present invention the interfering ribonucleic acid subject comprises two strands, whereby a 2'-O-methyl modified nucleotide and a non-modified nucleotide, preferably a nucleotide which is not 2'-O-methyl modified, are incorporated on both strands in an alternating fashion, resulting in a strand having a pattern MOMOMOMOM etc where M is a 2'-O-methyl nucleotide and O is a non-modified nucleotide. The same sequence of 2'-O-methyl modification and non-modification exists on the second strand, and preferably there is a phase shift between the two strands such that the 2'-O-methyl modified nucleotide on the first strand base pairs with a non-modified nucleotide(s) on the second strand and vice versa. This particular arrangement, i.e. base pairing of 2'-O-methyl modified and non-modified nucleotide(s) on both strands is particularly preferred in case of short interfering ribonucleic acids, i.e. short base paired double-stranded ribonucleic acids because it is assumed, although the present inventors do not wish to be bound by that theory, that a certain repulsion exists between two base-pairing 2'-O-methyl modified nucleotides which would destabilise a duplex, and particularly short duplexes, containing such pairings.

In a "phase shifted" arrangement of this nature, it is preferred if the antisense strand starts with a 2'-O-methyl modified nucleotide at the 5' end whereby consequently the second nucleotide is non-modified, the third, fifth, seventh and so on nucleotides are thus again 2'-O-methyl modified whereas the second, fourth, sixth, eighth and the like nucleotides are non-modified nucleotides. Again, not wishing to be bound by any theory, it seems that a particular importance may be ascribed to the second, and optionally fourth, sixth, eighth and/or similar position(s) at the 5' terminal end of the antisense strand which should not comprise any modification, whereas the most 5' terminal nucleotide, *i.e.* the first 5' terminal nucleotide of the antisense strand may exhibit such modification with any uneven positions such as first, optionally third, fifth and similar position(s) at the antisense strand may be modified. In further embodiments the modification and non-modification, respectively, of the

modified and non-modified nucleotide(s), respectively, may be any modification as described herein.

Although not limited thereto, the double-stranded structure of the inventive ribonucleic acid, which is also referred to as duplex, is formed by the first strand and second strand, respectively, or the first and second stretch of contiguous nucleotides. The length of the first stretch and second stretch, respectively, is typically about 15 to about 23, preferably 17 to 21, and more preferably 18 or 19 bases. A length of less than 30 nucleotides, preferably less than 21 nucleotides does not induce an interferon response in any biological system which is capable of showing RNA interference and an interferon response. This apparently is because a given cell is experiences profound physiological changes when double-stranded RNA longer than 30 base pairs binds and activates the protein kinase PKR and 2',5'-oligoadenylate synthetase. Activated PKR stalls translation via phosphorylation of eIF2a, activated 2',5'-AS causes mRNA degradation. These effects are not desired in target validation and animal models because they override the effect of the target specific knockdown on the phenotype.

RNAi oligonucleotides with loop structures

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In a sixth strategy for designing interfering ribonucleic acids of the present invention, the ribonucleic acid comprises a double-stranded structure where the double-stranded structure has a first strand comprising a first stretch of contiguous nucleotides that is at least partially complementary to a target nucleic acid, and a second strand comprising a second stretch of contiguous nucleotides that is at least partially identical to a target nucleic acid, and where one terminus of the first strand and one terminus of the second strand are linked by a loop structure.

In one embodiment the loop structure is comprised of a non-nucleic acid polymer. Suitable non-nucleic acid polymers include polyethylene glycol or similar polymers. The polymer may be chosen in such a manner that the two linked strands may hybridize to each other. This requires that the polymer has to have a certain molecular "hinge" structure or molecular flexibility to allow the bending of the molecule so as to allow that both stretches get in close contact and in a three-dimensional orientation which permits hybridization. In principle any molecule which complies with this requirement may be used in connection with the present invention. For example, amino acid based molecules may also be used.

Such amino acid based molecules may be either homopolymers or hetereopolymers. A useful example is a homopolymer consisting of seven glycine residues which allows the generation of a hinge as required to bring the two stretches to hybridize in the close proximity as needed. A suitable glycine based hinge is described, *e.g.*, in Guan K. L. and Dixon J. E. (1991), *Anal. Biochem.* 192, 262. In another embodiment the hinge may be formed by crown ethers of a type that is known in the art.

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In an alternative embodiment the loop is comprised of a nucleic acid moiety. In the context of these loops, LNA as described in Elayadi and Corey (2001) Curr. Opin. Investig. Drugs. 2(4):558-61 and Orum and Wengel (2001) 10 Curr. Opin. Mol. Ther. 3(3):239-43; and PNA are regarded as nucleic acids and may also be used as loop forming polymers. Basically, the 5'-terminus of the first strand may be linked to the 3'-terminus of the second strand. As an alternative, the 3'-end of the first strand may be linked to the 5'-terminus of the second strand. 15 The nucleotide sequence forming the loop structure is, in general, not critical. However, the length of the nucleotide sequence forming such loop seems to be critical for steric reasons. Accordingly, a minimum length of four nucleotides appears to be appropriate to form the required loop structure. In principle, the maximum number of nucleotides forming the hinge or the link between both 20 stretches to be hybridized is not limited. However, the longer a polynucleotide is, the more likely secondary and tertiary structures are formed and this can affect the required orientation of the stretches that must form the hybridised structure. Preferably, a maximum number of nucleotides forming the hinge is about 12 nucleotides. It is within the scope of the present invention that any of the designs 25 described above may be combined with this sixth strategy, i.e. by linking the two strands covalently in a manner that back folding (loop) can occur through a loop structure or similar structure.

The present inventors surprisingly have found that if the loop is placed 3' of the antisense strand, *i.e.* the first strand of the ribonucleic acid(s) according to the present invention, the resulting RNAi molecules have higher activity than molecules where the loop is placed 5' of the antisense strand. This result is contrary to the conventional wisdom that the orientation of the loop is irrelevant, based on the assumption that any RNAi, is subject to a processing during which non-loop linked RNAi is generated. If this was the case, however, the clearly

observed increased activity of those structures having the loop placed 3' of the antisense could not be explained. Accordingly, a preferred arrangement in $5' \rightarrow 3'$ direction of this kind of small interfering RNAi is second strand - loop - first strand.

Expression of RNAi from a vector

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The respective constructs may be incorporated into suitable vector systems. Preferably the vector comprises a promoter for the expression of RNAi. Suitable promoters include pol III and the U6, H1, 7SK promoters as described in Good et al. (1997) *Gene Ther*, 4, 45-54.

Use of RNAi for gene knockdown and target validation

The ribonucleic acid molecules according to the present invention have general applicability and permit knockdown or knockout of any desired coding nucleotide such as an mRNA, *i.e.* the expression of any gene producing an RNA may be modified. A particular application is the use of the inventive ribonucleic acid for target validation. As used herein, target validation means a process that involves determining whether a DNA, RNA, or protein molecule is directly involved in a biological process. Preferably the biological process is causally involved in a disease or non-standard condition and the gene under study is therefore a suitable target for development of a new therapeutic compound

Sequence homology studies have successfully classified genes into target families. The enormous task of deciphering which of these targets are key players in diseases and which should be used for subsequent drug development needs to be addressed in an efficient manner. Therefore, the knockdown of gene expression should be reduced by 50-100%, preferably by 90% to see significant effects on the phenotype. In other cases depending on the gene, a knockdown of as little as 20% might be sufficient to yield a phenotype. A phenotype will be defined by comparison of cells containing functional RNAi molecules with cells containing non functional RNAi molecules. This will ensure a significant readout even under conditions where the protein function is inhibited only partially. Generally there is no linear correlation between the degree of mRNA reduction and the extent of the change in phenotype. For some proteins a reduction of about 20 % of the protein is sufficient to create a change in the phenotype whereas in case of other genes and mRNA, respectively, as little as 5 to 10 % remaining protein is sufficient to maintain an observed phenotype.

A further use of the ribonucleic acid molecules according to the present invention is its use for the manufacture of a medicament or its use as a medicament. Such a medicament may either be used for the treatment and/or prevention of diseases or conditions such as any type of cancer where a gene and its product have been linked to the onset, cause or progression of this disease. Additionally, such a medicament may be used to treat diseases where the presence or overexpression of a gene product causes a pathological phenotype. In a preferred embodiment the disease is characterised by a gain of function and may be remedied through application or administration of the corresponding, biologically active RNAi. Diseases or conditions which may be treated by the medicament comprising a ribonucleic acid as disclosed herein, may be selected from the group comprising cancer, heart diseases, metabolic diseases, dermatological diseases, inflammatory diseases, immune system disorders and autoimmune disorders. The various forms of cancer include, but are not limited to, solid tumors and tumors of the hematopoietic system, such as glioblastoma, prostate cancer, breast cancer, lung cancer, liver cancer, pancreatic cancer and leukaemia. Metabolic diseases include, but are not limited to, obesity and diabetes. Dermatological diseases include, but are not limited to, psoriasis.

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In another aspect the ribonucleic acid molecules according to the present invention may be used as diagnostics, preferably for those diseases as specified in connection with the above-mentioned diseases and conditions. Such diagnosis could be based on the observation that upon applying the ribonucleic acid molecules according to the present invention to a sample which preferably contains cells, a change in the expression pattern of the sample occurs. Preferably such sample comprises cells from a subject from whom it is assumed that it may exhibit said disease or condition to be treated or a predisposition therefor.

A further application of the nucleic acids according to the present invention resides in their use in the screening of pharmaceutically active compounds and/or lead optimization. The latter is done such as to monitor or determine the effect of candidate drugs such as small molecules and compare the effect created by said candidate drugs with the effect observed upon administering specific RNAi designed on the basis of the principles disclosed herein. In doing so candidate drugs having off-target effects may be eliminated from the screening process whereas those candidate drugs which create a similar or identical

phenotype are deemed as highly relevant lead compound or may even be a pharmaceutically active compound themselves. In this approach, the highly specific RNAi molecules act as a gold standard against which candidate drugs are measured.

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In a further aspect the invention is related to a cell, preferably a knockdown cell, which contains a ribonucleic acid as disclosed herein. Such a cell preferably is either isolated or contained in a tissue or even organ which again preferably is not contained in an organism. However, the cell may also be contained in an organism. The cell is preferably a cell which is involved in the disease or condition which is to be treated by the ribonucleic acids of the invention. This kind of knock-down cells may be used to generate an expression profile based on, *e.g.*, mRNA or protein, in order to elucidate functional relationship and to determine downstream targets.

In a further aspect the invention is related to an organism containing a ribonucleic acid as disclosed herein. Preferably such organism is a vertebrate organism and more preferably the vertebrate organism is a mammal. A mammal as used herein is, among others and not limited thereto, an ape, a dog, a cat, a goat, a sheep, a pig, a guinea pig, a rabbit, a mouse, a rat and a human being.

In a still further aspect the present invention provides compositions containing one or more ribonucleic acids according to the present invention. Preferably such composition comprises negative and positive controls either in combination with the effective ribonucleic acid or separated therefrom. Such composition may further comprise a solvent, preferably a buffer.

In a further aspect the present invention is related to a pharmaceutical composition containing a ribonucleic acid according to the present invention and a pharmaceutically acceptable carrier. Pharmaceutically acceptable carriers are known in the art and comprise, among others, diluent, buffers and the like. Suitable carriers are described, for example, in Remington's Pharmaceutical Sciences (18th edition, 1990, Mack Publishing Co.). The pharmaceutical composition may comprise additional pharmaceutically active compounds. In cases where the disease or condition to be treated using the ribonucleic acid molecules of the present invention is the same or related disease of condition as that treated by the additional pharmaceutically active compound(s) then the

different mode of action of the ribonucleic acid molecules and the additional compounds will produce synergistic effects.

The invention is now further illustrated by reference to the figures and examples from which further features, embodiments and advantages of the present invention may be taken.

Example 1: Dose response of synthetic duplex RNAi molecules

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In this example the impact of NH₂ end protection groups on the activity of duplex RNAi molecules was investigated. Synthetic siRNAs were purchased from Biospring (Frankfurt, Germany). The ribo-oligonucleotides were resuspended in RNase free TE to a final concentration of 50 μM. In the case of bimolecular siRNA molecules equal aliquots (100 μM) were combined to a final concentration of 50 μM. For the formation of intramolecular duplexes the siRNAs were incubated at 50 °C for 2 min in annealing buffer (25mM NaCl; 5mM MgCl₂) and were cooled down to RT. Transfections were carried out in 96 well or 10-cm plates (at 30% to 50% confluency) by using various cationic lipids such as Oligofectamine, Lipofectamine (Life Technologies), NC388 (Ribozyme Pharmaceuticals, Inc., Boulder, CO), or FuGene 6 (Roche) according to the manufacturer's instructions. RNAi molecules were transfected by adding preformed 5x concentrated complex of annealed RNAi and lipid in serum-free medium to cells in complete medium. Prior to transfection 2500 HeLa cells were plated per well 15 – 18 hours before transfection for the 96 well format.

The total transfection volume was 100 μ l for cells plated in 96-wells and 10 ml for cells in 10 cm plates. The final lipid concentration was 0.8 to 1.2 μ g/ml depending on cell density; the RNAi concentration is indicated in each experiment.

Complexes were added to cells to yield a final 1x concentration of both lipid and RNAi. Depending on the analysis performed following transfection, cells were lysed using a standard cell lysis buffer for protein extraction (Klippel, A., Escobedo, J. A., Hirano, M. and Williams, L. T. (1994). Mol Cell Biol 14, 2675-2685) or a denaturing buffer for RNA isolation according to the RNA isolation kit

(Invitek, Berlin (Germany) 24 to 48 hours post transfection for RNA analysis and 48 to 72 hours post transfection for protein analysis by Western Blot.

Determination of the relative amounts of RNA levels by Taqman analysis:

5 24h post transfection the RNA of cells transfected in 96-wells was isolated and purified using the Invisorb RNA HTS 96 kit (InVitek GmbH, Berlin). Inhibition of PTEN mRNA expression was detected by real time RT-PCR (Taqman) analysis using 300 nM PTEN 5' primer CACCGCCAAATTTAACTGCAGA, 300 nM PTEN 3' primer 10 AAGGGTTTGATAAGTTCTAGCTGT and 100 nM of the PTEN Tagman probe Fam-TGCACAGTATCCTTTTGAAGACCATAACCCA-Tamra in combination with 40 nM β-actin 5' primer GTTTGAGACCTTCAACACCCCA, 40 nM βactin 3' primer GACCAGAGGCATACAGGGACA and 100 nM of the β-actin Taqman probe Vic-CCATGTACGTAGCCATCCAGGCTGTG-Tamra. The Akt 15 primers and probes are determined in Sternberger et al. (Sternberger, supra) and are used according to the manufacturer's instructions (Applied Biosystem; use of Amplicon Set). Also said primers and probes may be designed using the software program Primer Express (Applied Biosystem). The reaction was carried out in 50 μl and assayed on the ABI PRISM 7700 Sequence detector (Applied Biosystems) 20 according to the manufacturer's instructions under the following conditions: 48°C for 30 min, 95°C for 10 min, followed by 40 cycles of 15 sec at 95°C and 1 min at 60°C.

RNA knockdown was shown by real time RT-PCR analysis of HeLa cells transfected with 21 nt long siRNA duplex molecules unmodified and modified with either NH $_2$ or inverted Abasics groups at the 5'-end at a lipid carrier concentration of 1.0 μ g/ml. Cell density was 2000 cells/well. Modifications on the 3'-end were either RNA overhangs, RNA overhangs with amino groups or DNA overhangs.

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Preparation of cell extracts and immunoblotting. Cells were washed twice with cold phosphate-buffered saline and lysed at 4°C in lysis buffer containing 20 mM Tris (pH 7.5), 137 mM NaCl, 15% (vol/vol) glycerol, 1% (vol/vol) Nonidet P-40 (NP-40), 2 mM phenylmethylsulfonyl fluoride, 10 mg aprotinin per ml, 20 mM leupeptin, 2 mM benzamidine, 1 mM sodium vanadate, 25 mM β-glycerol phosphate, 50 mM NaF and 10 mM NaPPi. Lysates were

cleared by centrifugation at 14,000 x g for 5 minutes and aliquots of the cell extracts containing equal amounts of protein were analyzed for protein expression by Western-blotting: Samples were separated by SDS-PAGE and transferred to nitrocellulose-filters (Schleicher & Schuell). Filters were blocked in TBST buffer (10 mM Tris-HCl (pH 7.5), 150 mM NaCl, 0.05% (vol/vol) Tween 20, 0.5% (wt/vol) sodium azide) containing 5% (wt/vol) dried milk. The respective antibodies were added in TBST at appropriate dilutions. Bound antibody was detected using anti-mouse- or anti-rabbit-conjugated horse radish peroxidase (Transduction Laboratories) in TBST, washed, and developed using the SuperSignal West Dura (Pierce) or ECL (Amersham) chemoluminescence substrates (c.f. Sternbergeret al. (2002). Antisense. Nucl. Ac. Drug Dev. in press..

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Antibodies. The murine monoclonal anti-p110 antibody U3A and the murine monoclonal anti-p85 antibody N7B have been described (Klippel et al., 1994, supra). Rabbit polyclonal anti-Akt and anti-phospho Akt (S473) antibodies were obtained from Cell Signaling Technology. The murine monoclonal anti-PTEN antibody was from Santa Cruz Biotechnology. The PTEN 53 specific antisense molecule, *i.e.* geneBloc, is described in Sternberger *et al.* [Sternberger, *supra*] having the following sequence (ucuccuuTTGTTTCTGcuaacga), whereby the nucleotide depicted in lower case are ribonucleotides whereas the nucleotide in capital letters are desoxyribonucleotides. This antisense molecule is also identical to RNAi 1A without TT.

The results are shown in Fig. 3A and the respective RNAi molecules in Fig. 3B which are directed to the mRNA of PTEN. The nucleotides written in lower case letters represent ribonucleotides whereas capital letters represent desoxyribonucleotides. The term NH₂ indicates that the 3' '-position of the ribonucleotide was modified by an amino group. The RNAi molecules used in this and other examples disclosed herein are also referred to as small interfering RNA molecules siRNA. It is to be noted that in any of the figures contained herein the upper strand is the antisense or first strand, whereas the lower strand is the sense or second strand of the interfering RNA molecule.

As can be taken from Fig. 3A amino end modifications such as amino modification and inverted abasics modification of the terminal OH group of the nucleic acid are as potent as unmodified ends when the modification is located at the 3' end of the antisense strand (see also Fig. 8A; 8B). Therefore chemical

modification to stabilize or with other beneficial properties (delivery) will be tolerated without activity loss when located at the 3' OH; especially when the 3'OH is located on an overhanging nucleotide.

For the experiment shown in Fig. 3C similar conditions as outlined above were used. The first strand and the second strand of the RNAi were either modified by a NH₂ group at the 3'-position of the ribose moiety or by an inverted abasic at said positions. The first construct is designated as siRNA-NH₂ (3A3B) the second as siRNA-iB (4A4B). The sequence of both molecules is depicted in Fig. 3B. The term 3A3B indicates that the interfering ribonucleic acid consists of strand 3A as the antisense strand and strand 3B as the sense strand. For reason of comparison an antisense oligonucleotide designated GB53 (Sternberger et al., supra) was generated which was directed against the PTEN mRNA as well. The particularities of this latter experiment were as follows.

As may be taken from Fig. 3C end protected RNAi molecules depicted in Fig. 3B are functional in yielding a PTEN protein knockdown.

From this example it can be taken that both end protection groups render RNAi molecules active in knocking down PTEN protein. This inhibition is as efficient as inhibition with antisense constructs but at lower concentrations used which is a clear advantage over the already very powerful antisense technology.

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Example 2: Overhang requirements for RNAi duplex activity in vivo

The experimental procedures were the same as depicted in connection with Example 1 except that the PTEN mRNA targeting interfering RNAi molecules were differently designed. The results are shown in Fig. 4A as dose response curves with Fig. 4B showing the particular sequence and modifications of the interfering RNAi molecules used to generate the data depicted in Fig. 4A. The nomenclature is such that, *e.g.*, RNAi 18 is composed of strand 18A as antisense strand and strand 18B as sense strand.

Blunt ended molecules were compared to molecules with 3'-overhangs (RNAi 18) and 5'-overhangs (RNAi 30 and RNAi 31) in their activity to knockdown PTEN mRNA in HeLa cells. The activity of blunt ended molecules (RNAi 28) and molecules with 5'-overhangs was comparable to the activity of molecules with 3'-overhangs. This shows that 3'-overhangs are not required for RNAi activity.

Example 3: Duplex length requirements of interfering RNA molecules for RNAi activity in vivo

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The experimental approach was similar to the one outlined in connection with Example 1 except that the interfering RNA molecules were directed against the mRNA of Akt1. The negative control to show the specificity of the RNAi molecules was again p110 mRNA. The experimental results are shown in Fig. 5A with the particularities of the interfering RNAi molecules used being represented in Fig. 5B. Similar experiments were carried out with further siRNA constructs which are depicted in Fig. 7A, left panel, whereby the arrows indicate mismatches and desoxyribonucleotides are expressed in capital letters. The inhibition of Akt1 mRNA expression in HeLa cells transfected with the indicated amounts of siRNA molecules is depicted on the right panel of Fig. 7A.

Taqman analysis on Akt RNA from HeLa cells transfected with different RNAi molecules shows that the doublestrand duplex of the siRNA molecules has to be longer than 17 base pairs to show activity whereas molecules with 17 base pair long duplexes or shorter are not functional even if sequence-specific overhangs are added. The shortest RNAi molecules successfully tested were 18 to 19 nucleotides or base pairs in length. It is to be noted that the design of the interfering RNA molecule 51A/51B referred to as RNAi 51 corresponds to the one as described in international patent application WO 01/75164. The RNAi molecule 55A/55B comprises a stretch of 17 nucleotides and has a clearly decreased activity in terms of degradation of Akt1 mRNA.

As may be taken from Fig. 7A 19 nt long duplexes are highly efficient in reducing Akt1 mRNA levels independent of the nature (desoxy- or ribonucleotides) of the 3' overhang (compare molecules 1AB, 2AB, 3AB, 4AB). The 17 nucleotide long siRNA (molecule 5AB) showed a dramatically reduced silencing activity confirming the above expressed understanding that active siRNA duplexes should be at least 18 nt or longer. Without wishing to be bound by any theory this result may be explained mechanistically by two different requirements. First, a minimum base pairing of 18 nt between the antisense of the siRNA and the target mRNA may be obligatory, or second, incorporation into the RNA-induced silencing complex (RISC) requires a minimum length of the siRNA duplex. To address this question a 19 nt long siRNA duplex molecule with one

and two terminal mutations (CG and UA inversion) relative to the wild type sequence was synthesised (molecules 6AB and 7AB). Both molecules, even the molecule with a stretch of only 15 nt base pairing to the target mRNA were functional in inducing the Akt1 mRNA level. Therefore, it can be concluded that the duplex length itself, but not the base pairing of the antisense siRNA with the target mRNA seems to determine the minimum length of functional siRNAs. This suggests that the length of the double-stranded helix is an important determinant for the incorporation into the RISC complex. The introduced mismatches at the terminal ends of the siRNA duplexes had little effect on RNA interference.

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Given the experimental results, the minimum requirement for optimum RNAi mediated interference is thus a duplex length of 18 or 19 nucleotides, independent of the further design of the RNAi molecules such as blunt end or 5'-overhang or any other form as disclosed herein, but generally applicable to RNAi molecules. However, it has to be acknowledged that the particular design of the RNAi molecules may confer further advantages to said molecules, such as, *e.g.*, increased efficiency and increased stability, respectively.

Example 4: Target-antisense homology requirements for RNAi in vivo

The experimental set-up was similar to the one described in example 1, whereby the RNAi is specific for Akt1. In addition, a PTEN specific interfering RNA molecule was designed and used as negative control. The results are shown in Fig. 6A and Fig. 6B. Basically the same experiment was carried out using further siRNA molecules as depicted in Fig. 7B with the results being indicated in Fig. 7B (right panel) and Fig. 7C, respectively.

Having established the minimal duplex length of 18 or more than 18 nucleotides for functional siRNA molecules we asked the question how many matching nucleotides between target mRNA and siRNA are necessary for silencing activity. As shown by Taqman analysis on Akt1 RNA a stretch of 19 to 15 nucleotides perfectly matching to the target RNA, in the case of Akt 1, was sufficient to mediate RNAi activity. A PTEN specific RNAi molecule did not reduce RNA amounts of Akt1 thus confirming the specificity of this approach. Molecules containing mismatches of one or two nucleotides at any or both ends of a strand were functional suggesting that a homolog stretch of 15 nt between a target mRNA and RNAi is sufficient for gene silencing.

It can be concluded from these data that unspecific gene silencing can occur by chance through unspecific binding to unrelated targets. This is based on the understanding that a stretch of 15 to 17 matching base pairs is not specific for a single gene and will occur by chance considering the complexity and size of the genome or transcriptosome of vertebrates. Apart from the above mentioned experiments the location of the mismatch also was subsequently analysed. For this purpose a 19 nt long blunt siRNA directed against PTEN mRNA was used. The sequence changes in one siRNA strand were compensated by complementary changes in the other strand to avoid disrupting duplex formation. As may be seen from both Figs. 7B and C, respectively, a siRNA with only one point mutation in the centre of the molecule was severely compromised in its ability to use mRNA and protein expression levels. This result indicates that the RNA machinery is highly discriminative between perfect and imperfect base pairing between target mRNA and siRNA in the centre of the duplex. This extreme dependence on a perfect complementarity between target and siRNA has already been described for RNAi interference in the Drosophila system, however, not yet in connection with mammalian systems such as HeLa.

Based on this observation the present invention reduces this off-target problem of siRNA by two approaches. First by reducing the molecule length of the siRNA molecules to the minimal requirements (18-19 nt) and thereby reducing the chance of homology to off-targets. Second, by inactivation of the sense strand to prevent a unwanted RNA silencing caused by accidental complementarity of the sense strand to a unrelated target RNA (see also Example 6).

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Example 5: Stability of modified RNAi molecules in serum

Oligonucleotides were incubated in human serum for 15 min and two hours and loaded on 10% polyacrylamide gel with untreated controls. The results are shown in Fig. 8A. The various RNAi molecules used are shown and described in more detail in Fig. 8B.

From this example it can be taken that the RNAi duplex of RNA molecules with all nucleotides modified with 2'-O-methyl groups (RNAi molecules 79A79B and 28A28B) have higher stability in serum. It is also shown that a blunt duplex is more stable than the duplex molecule with overhangs. From

this the conclusion may be drawn that end protection (e.g. iB or amino) does not increase the stability in serum. In addition, it can also be concluded that, in contrast to the understanding in the art before the filing of the present application, endonucleases are more important than exonucleases in the protection of RNAi molecules.

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In view of this, in addition to the various modifications or designs of the inventive RNAi molecules as disclosed above, further or additional modification of the nucleotides may include the use of a phosphorothioate backbone of the RNAi molecules which may be either complete or partial in order to inhibit endonuclease function. A complete phosphorothioate backbone means that any of the nucleotides exhibits a phosphorothioate group whereas a partial phosphorothioate backbone means that not all of the nucleotides forming the RNAi molecule have a phosphorothioate modification. This modification is suitable to increase the lifetime of RNAi molecules irrespective of the further design of RNAi molecules. In this regard, the present invention encompasses a partially or completely phosphorothioate modified RNAi which may be realized in connection with the different strategies for the design of interfering RNA molecules as disclosed herein or with any of the designs known in the art.

20 Example 6: Inactivation of the sense strand by NH₂ end protection groups on the 5' and 3' ends

The experimental set-up was similar to the one described in connection with Example 1 with the target nucleic acid sequence being PTEN mRNA. The concentration of HeLa cells was 2,000 cells per well. RNA of PTEN was analysed in Taqman assays after transfection of differently modified RNAi molecules. The different interfering RNA molecules used are depicted in Fig. 9B and the experimental results are shown in Fig. 9A.

The dose response curves of various RNAi molecules depicted in Fig. 8A show that RNAi molecules are functional when the sense strand, *i.e.* the second stand, is modified on both ends with amino groups. Particularly effective are RNAi molecules 20A26B, 18A26B, and 28A26B. The lowest activity is shown by RNAi molecule 26A26B which corresponds to end modification on all 4 ends of the duplex (the molecules described by Tuschl are 18AB).

However, RNAi activity is also achieved when the antisense strand, *i.e.* the first strand, is modified only at the 3' end leaving a free OH group at the 5' end (RNAi constructs 22A26B; 20A26B). There is no activity when the antisense strand is modified with amino groups on both the 5' and the 3' end (26A26B).

5 This leads to the conclusion that any end of the antisense strand and more particularly the 5' end of the antisense should be kept without modifications. Additionally, it is worth stating that the NH₂ end modification can be used to inactivate the sense strand on the 5' and 3' end and therefore reduce off-target effects mediated by an otherwise functional sense strand which results in a significantly increased specificity of the RNAi molecule which is advantageous for target validation as well as for any medical use of the RNAi molecule.

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The further generalisation of the results from this experiment is depicted in Fig. 9C. Functionally active RNAi are accordingly those not having an amino modification at the antisense strand or having an amino modification only at the 3' end of the antisense strand whereas an amino modification at both ends of the antisense strand is not functional, *i.e.* does not result in a knockdown of the target mRNA.

Example 7: Impact of 2'-O-methyl modification of RNAi molecules for endonuclease protection.

RNA knockdown was again shown using real time RT-PCR analysis on HeLa cells transfected with RNAi duplex molecules directed against the PTEN mRNA as represented in Fig. 10A. Experimental procedures were basically the same as specified in Example 1. The structure of the RNAi molecules investigated and their dose responses, which are depicted in Fig. 10A, are shown in Fig. 10C. The nucleotides printed in bold are those having a 2'-O-methyl modification.

It is illustrated by the dose response curves shown for various RNAi molecules in Fig. 10A that internal 2'-O-alkyl groups reduce RNAi activity. Preferably such 2'-O-alkyl groups are 2'-O-methyl or 2'-O-ethyl groups. However, molecules with unmodified nucleotides in combination with 2'-O-alkyl modification show significant activity. As is also depicted in Fig. 10A no activity was obtained when the antisense strand is all modified with 2'-O-methyl groups and the sense strand is not modified (*c.f.*, *e.g.*, RNAi molecule 79A28B). Taken the results of a stability test such as incubation of the various RNAi molecules in

serum, as depicted in Fig. 10B, shows that 2'-O-alkyl modifications stabilize RNAi molecules against degradation. This clearly beneficial effect, however, is at least to a certain degree counterbalanced by the effect that 2'-O-alkyl modifications generally result in a reduced knockdown activity. Accordingly, the design of RNAi molecules has to balance stability against activity which makes it important to be aware of the various design principles as disclosed in the present application.

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Example 8: Impact of blocks of internal 2'-O-methyl modifications on the stability of RNAi molecules in serum.

The experimental approach in connection with this study was the same as depicted in Example 1. Again, PTEN RNA was analysed by real time RT-PCR on HeLa cells at a density of 2000 cells/well which were transfected with different doses of RNAi molecules. RNAi molecules were incubated in serum for two hours and analysed on a 10% polyacrylamide gel The results of this study are illustrated in Figs. 11A to 11C, whereby Fig. 11A shows the dose response of the various RNAi molecules depicted in Fig. 11C and Fig. 11B shows the result of a stability test using some of the RNAi molecules depicted in Fig. 11C. The nucleotides written in bold in Fig. 11C are the ones carrying a modification, in this case a 2'-O-methyl modification of the ribose moiety of the nucleotides.

A dose dependent inhibition by the unmodified RNAi molecules was observed. It was also shown that the 2'-O-methyl modification of the core 9 nucleotides made the RNAi stable in serum and allowed activity of the duplex in mediating the interference phenomenon leading to a degradation of the PTEN mRNA. Total modification of the sense strand makes the RNAi molecule stable in serum and allows certain activity.

Alternating blocks of 5 nucleotides with 2'-O-methyl modification renders the RNAi molecule stable in serum and allows activity on PTEN RNA as shown by incubating the RNAi duplex in serum for two hours and loading the samples on a 10% polyacrylamide gel. As may be taken from Fig. 11B the duplex comprising strands 80A and 80B was strongly degraded after incubation in serum for two hours. The duplex consisting of strands 82A and 82B confirmed the result that the 5'-end of the first strand which comprises the antisense strand should not be modified at the 5'-terminal nucleotides (compare 82A82B with the reverse

orientated 81A81B). This was also confirmed by the results obtained having the duplex consisting of the strands 86A and 86B which is both active and stabilised in serum. It is noteworthy that molecules with unmodified blocks at the terminal 5' of the antisense strand are more active whereby the 5'terminal OH group is preferably not derivatized.

Further experiments were carried out using different modification patterns of 2'O-methyl modification of the nucleotides. The results thereof are shown in Fig. 12A to 12C and further discussed herein in example 9.

10 Example 9: The impact of alternating internal 2'-O-alkyl modification on serum stability of RNAi molecules.

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The experimental set-up for performing this kind of study was the same as used in connection with the studies reported in Example 1 and Example 8, respectively, with the targeted nucleic acid being again PTEN mRNA. HeLa cells were transfected with the different RNAi molecules depicted in Fig. 12B and RNA knockdown was demonstrated using real time RT-PCR on PTEN RNA in a dose-dependent manner (Fig. 12A). The stability of the various RNAi molecules after 15 min and two hours in serum at 37°C is depicted in Fig. 12C and a Western Blot for p110 and PTEN as the target-protein of the various RNAi molecules is depicted in Fig. 12D with the RNAi molecules tested being the same in both the experiments underlying Fig. 12C and Fig. 12D.

As illustrated in Fig. 12A and Fig. 12C nucleotides modified with 2'-O-methyl groups alternating with unmodified nucleotides rendered RNAi molecules stable in serum while still allowing them to be active in the sense of interfering with the target mRNA. It was shown that incubation of RNAi duplex molecules for 15 min and two hours in serum degraded the unmodified duplex and the duplex where the 10 most 5'-positioned nucleotides were unmodified.

In the RNAi molecules represented in Fig. 12B various patterns of modified and unmodified nucleotides were prepared. The RNAi molecule 94A1/94B1 comprises a structure wherein a modified nucleotide is flanked by an unmodified nucleotide with the unmodified nucleotide being located at the 5' end of the first strand. The RNAi molecule comprised of strands 94A2 and 94B2 was another example where the modified nucleotides and the unmodified nucleotides of the first and the second strand were located at opposing sites. In contrast to this

the RNAi molecule comprised of strands 94A1 and 94B2 had the same pattern of modified and unmodified nucleotides. However, there was a phase shift such that the modified nucleotide base paired with an unmodified nucleotide. The two RNAi molecules comprised of strands 94A1 and 94B1 and strands 94A2 and 94B2 differed from each other such that in the first case the first strand starts with an unmodified nucleotide and the corresponding first nucleotide of the second strand, *i.e.* the nucleotide at the 3' end of the second strand, starts with an unmodified nucleotide with the arrangement being opposite to this in the RNAi molecule comprised of 94A2 and 94B2.

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Additionally, alternatingly modified RNAi molecules as depicted in Fig. 12B were functional in mediating a PTEN protein knock down as shown in Fig. 12D but only when the second 5' and second 3' terminal nucleotide was not modified (see 94A294B1 and 94A294B2). Taken together these data show that the most stable and most active RNAi molecules have alternating 2' alkyl modified and unmodified nucleotide residues. It should be noted that these molecules do show a very similar mRNA reduction when compared to unmodified siRNA molecules while being stable in serum and therefore allowing increased or easier handling.

20 Example 10: Functional Protein Knockdown mediated by internally modified RNAi molecules

The experimental approach was similar to the one outlined in connection with example 1.

Western Blots were performed on HeLa cells harvested at various time points following transfection (48, 72, 96 and 120 hours) with alternatingly modified RNAi molecules as depicted in Fig. 12B. For experimental reasons it is noteworthy that at the 96 hour time point cells were split and half the population was replated. A total of 40 nM of the various RNAi molecules were applied to the cells. Cells were continuously transfected for 72h with cationic lipids as described in example 1; then replated in the absence of transfection reagents.

Transfections were carried out in 96 well or 10-cm plates (at 30% to 50% confluency) by using various cationic lipids such as Oligofectamine, Lipofectamine (Life Technologies), NC388, L8 (Atugen, Berlin), RNAi were transfected by adding pre-formed 5x concentrated complex of siRNAs and lipid in

serum-free medium to cells in complete medium. The total transfection volume was 100 μ l for cells plated in 96-wells and 10 ml for cells in 10 cm plates. The final lipid concentration was 0.8 to 1.2 μ g/ml depending on cell density; the siRNA concentration is indicated in each experiment.

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The result of the Western Blot analysis is depicted in Fig. 13. As can be taken from this Figure, modified RNAi molecules of the 94A2B1 and 94A2B2 version yielded a longer lasting PTEN protein knock down as unmodified molecules. The lack of protein knock down also seen in Fig. 12 with molecules of the 94A1B1 and 94A1B2 version was confirmed in this experiment. Unmodified molecules (80AB) were not as potent in supporting long lasting protein knock down when the cells were not continuously transfected.

Example 11: Persistent PTEN protein knock down with alternating 2'-O-methyl modifications of RNAi molecules

The experimental approach was similar to the one outlined in connection with example 10 with the exception that the transfection was terminated after 5 h by replacing the transfection medium with new medium. The protocol was slightly modified such that for each of the RNAi molecules a 40 nM concentration was realized using a stock solution of 1 µg RNAi / ml cationic lipid as described in connection with example 1.5 hours after transfection the medium was withdrawn and fresh EMEM added. The cells were split after 72, with half of the cells being lysed and the other half newly plated and lysed 24 h later (96h post transfection). The result of a Western Blot analysis using 3 different RNAi molecules (80AB, 94A1/B2, 94A2/B1 are depicted in Fig. 14. As a positive control untreated cells were used. Fig. 14 shows the expression of PTEN after 72 h and 96 h, respectively. Taken the structural particularities of the various RNAi molecules it can be taken from Fig. 14 that protein knockdown is persistent with alternating molecules of the 94A2B1 kind even over 96 h after splitting and replating cells compared to unmodified RNAi molecules (such as 80AB) and RNAi molecule 94A1B2.

A further experiment was carried out using the siRNA constructs as depicted in Fig. 15A (left panel). From the results as depicted as ratio of PTEN/p110α mRNA degradation at the various concentrations of siRNA constructs administered to the test system, it can be taken that siRNA molecules

with either one or both strands consisting of 2'-O-methyl residues were not able to induce RNA interference in the mammalian system (Fig. 15A, molecules V2, V5, V6). However, the decrease in activity was less pronounced when only parts of the strands were modified. Interestingly, a molecule having an unmodified antisense strand (which is the upper strand in the representation throughout this specification if not indicated otherwise) and a completely modified sense strand was significantly more active when compared to the reversed version (Fig. 5A, molecules V5 and V6). This result suggests that the antisense strand of the siRNA seems to be more critical and sensitive to modification. The most efficient molecules inducing PTEN mRNA had only stretches of modifications leaving the 5' end unmodified or were modified on alternating positions on both strands (Fig. 15A, molecules V10, V12).

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To test the nuclease resistance the different siRNA versions were incubated in serum followed by PAA gel electrophoresis. The result is shown in 15 Fig. 15B (right panel with the various sequences indicated on the left panel of Fig. 15B). As shown earlier, blunt ended siRNA molecules with unmodified ribonucleotides were very rapidly degraded whereas a complete substitution with 2'-O-methyl nucleotides mediated resistance against serum-derived nucleases (Fig. 15B, compare molecule AB with V1). siRNA molecules with partial 2'-O-20 methyl modification also showed an increased stability when compared to unmodified siRNAs. In particular, molecules with alternating modifications on both strands showed a significant improve in instability (Fig. 15B, molecules V13, V14, V15 and V12). More importantly, transfection of three of these molecules into HeLa cells resulted in a significant down regulation of PTEN protein 25 expression as depicted in Fig. 15C, length 6, 9 and 10). In this RNA interference activity assay an unexpected preference for molecules was observed which were modified at every second nucleotide beginning with the most 5' terminal nucleotide of the antisense strand (molecules V15 and V12). Molecules which contained modifications beginning with the second nucleotide at the 5' end of the 30 antisense strand were more stable but had a strongly reduced activity in gene silencing (molecules V13, V14). This result points towards highly specific interactions between the involved enzymes and precise nucleotides in the siRNA duplex. Taken together the data shown herein demonstrate that 2'-O-methyl

modifications at particularly selected positions in the siRNA duplex can increase nuclease resistance and do not necessarily abolish RNAi completely.

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Although an increased stability of synthetic siRNA has primarily implication for *in vivo* application, it was also analysed whether the particular modification can also lead to an extended protein knock-down in cell culture systems. Accordingly, HeLa cells were transiently transfected for six hours using different versions of PTEN specific siRNAs. The lipid siRNA complex was then washed away and the PTEN protein knock-down was analysed 48 hours and 120 hours later. Although knock-down experiments without continued transfection of siRNAs were complicated due to rapid growth of untransfected cells in this time period resulting in a very transient knock-down, the present inventors were able to demonstrate a prolonged PTEN protein knock-down with siRNA molecules stabilised by the described 2'-O-methyl modification. At 48 hours post transfection the unmodified siRNA (AB) showed the biggest reduction in PTEN protein levels, however, at 120 hours post transfection the reduction in PTEN protein expression was superior with the siRNA stabilised by alternating 2'-O-methyl modifications (Fig. 15D, compare lane 2 with lanes 4, 6 and 7).

From the results it can also be surmised that the starting nucleotide position of the alternating modification can affect the activity of the duplex. To test this preference in more detail two additional series of siRNAs were synthesized, one specific for the kinase Akt1 and the other one specific for p110\beta which is one of the two catalytic subunits of PI(3-) kinase. The particular constructs are shown in Fig. 16A. Only 19 nt long siRNA duplexes either without any or with 2'-O-methyl modification on every second other nucleotide were used. Using Akt1 as a target an efficient protein knock-down as well as a dramatic reduction in phospho-Akt levels was observed with blunt, unmodified siRNAs (Fig. 16A, right panel). From the different versions of molecules with modifications on every second other nucleotide only one was efficiently mediating RNAi (Fig. 16A, molecule V5). This siRNA molecule contained an antisense strand which was modified at the most terminal 5' and 3' nucleotides. The sense strand started with the unmodified nucleotides at the most terminal position, resulting in a structure in which the modified and unmodified ribonucleotides of both strands face each other. As expected, this molecule was

also protected against serum-derived nucleases as depicted in Fig. 16B (molecule V5).

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Interestingly, a very similar 19 nt long siRNA molecule (V4) with modifications beginning at the second nucleotide of the antisense strand showed no RNA interference activation in the particular assay used. The version V6 in which the modified nucleotides of the antisense strand face modified nucleotides on the sense strand, was also inactive in this experiment. An identical series of 19 nt long siRNA molecules specific for p110ß confirmed these observations as depicted in Fig. 16C. Again the similarily modified siRNA molecule (V5) was the most active, as indicated by reducing Akt phosphorylation, which is indicative for a reduced P (I)-3 kinase activity due to reduced p110β levels. The reduced activity of the molecules V6 might be explained by reduced duplex stability since the same structure was active in the PTEN knock-down experiment with 21mer siRNAs. Although it is known that 2'-O-methyl modification on both strands facing each other reduces the stability of nucleic acid duplexes, the difference between the activity of siRNA molecules V4 and V5 (Figs. 16B and C) probably is not due to differences in duplex stability since the number of base pairing of modified and unmodified nucleotides is identical. This difference in activity might be due to specific requirements in the interacting proteins involved in the degradation of the target mRNA. Also it can be taken from these experiments that the most terminal nucleotides of the antisense strand can be modified at the 2'-OH-group with significant loss of silencing activity.

Example 12: Different loop structures are functional in mediating RNA interference

To test whether RNAi molecules, preferably synthetic RNAi molecules with self complementary structures, can inhibit gene expression as efficiently as standard double stranded siRNA molecules, HeLa cells were transfected with p110β specific synthetic siRNAs. Transfections were carried out in 96 well or 10-cm plates (at 30% to 50% confluency) by using various cationic lipids such as Oligofectamine, Lipofectamine (Life Technologies), GeneBlocs were transfected by adding pre-formed 5x concentrated complex of GB and lipid in serum-free medium to cells in complete medium. The total transfection volume was 100 μl for cells plated in 96-wells and 10 ml for cells in 10 cm plates. The final lipid

concentration was 0.8 to 1.2 µg/ml depending on cell density; the RNAi molecule concentration is indicated in each experiment.

A dose dependent titration showed no significant difference in efficiency of mRNA knock-down achieved by the standard bimolecular double strand 21mer and the corresponding monomolecular molecules as analysed by realtime PCR (Taqman) (Fig. 17A). Two different loop structures a (A)₁₂ loop and a HIV derived pA-loop were tested in parallel with similar results. A comparision of the relative position of the antisense sequence and the loop structure revealed an improved knock-down efficiency with the antisense sequence being located 3' to the loop (Fig. 17B; compare construct 3A, 3B and 4A, 4B).

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Example 13: Studies on the efficacy of different intramolecular hairpin loops and intermolecular "bubbles"

The influence of different loop structures on inhibition of mRNA and protein expression was tested. For these experiments Akt1 and Akt2 were chosen as targets. The experimental approach was similar to the one described in example 12.

Significantly, the reduction in Akt1 mRNA as depicted in Fig. 18A and Fig. 18B as well as Akt1 protein levels as depicted in Fig. 18C was completely independent of the loop structure tested (compare molecules 5A, 6A, 7A, 8A) (the structure of the RNAi molecule tested is always depicted underneath the bar diagram). Even a molecule containing a rather unphysiological structure such as a polyethyleneglycol linker (PEG) as a loop efficiently reduced Akt1 expression indicating that size and nucleotide sequence of the loop is not crucial (Fig. 18A; molecule 8A). A synthetic siRNA molecule specific for Akt2 (9A) was used to control for specificity, and had no effect on Akt1 levels as shown in Fig. 15A. This molecule, however, efficiently silenced Akt2 expression (Fig. 18B; Fig. 18C). Self-complementary RNA molecules with loop structures have the possibility to anneal as double strands in monomolecular or bimolecular structures under physiological hybridization conditions (Fig. 18B, loop or bubble structure). To address the question of whether the siRNA molecules exert their function via adapting an intra-molecular loop or an inter-molecular "bubble" (schematically shown in Fig. 18B) two molecules not capable of folding back on themselves were transfected. These constructs contained Akt1- and Akt2-specific sequences

within the same molecule (Fig. 18B, constructs 10A, 10B) and were designed so as to prevent formation of an intramolecular duplex (*i.e.* a bimolecular duplex or "bubble" was formed). Surprisingly, this molecule efficiently mediated both Akt1 and Akt2 mRNA knock-down as well as protein knock-down when transfected after annealing of both strands.

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Whether loop and bubble structures are indeed substrates for RNA processing enzymes, e.g. Dicer, is not clear at this point. A recent study by Paddison and coworkers suggests that hairpin containing siRNAs are more dependent on Dicer activity than double stranded siRNAs. However, these data demonstrating RNA interference activity using a PEG linker molecule indicate that the linker sequence is likely to be irrelevant.

The features of the present invention disclosed in the specification, the claims and/or the drawings may both separately and in any combination thereof be material for realizing the invention in various forms thereof.